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- (71) Applicant (for all designated States except AT, US): NO-VARTIS AG [CH/CH]; Lichtstrasse 35, CH-4056 Basel (CH).
- (71) Applicant (for AT only): NOVARTIS-ERFINDUNGEN VERWALTUNGSGESELLSCHAFT M.B.H. [AT/AT]; Brunner Strasse 59, A-1230 Vienna (AT).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): BAIR, Kenneth, Walter [US/US]; 95 Melrose Road, Mountain Lakes, NJ 07046 (US). GREEN, Michael, A. [US/US]; 2180 Biddle Lane, Easton, PA 18040 (US). PEREZ, Lawrence, B. [US/US]; 12 Windsor Place, Hackettstown, NJ 07840 (US). REMISZEWSKI, Stacy, W. [US/US]; 147 Honeysuckle Drive, Washington Township, NJ 07676 (US). SAMBUCETTI, Lidia [US/US]; 32 Lone Mountain

Court, Pacifica, CA 94044 (US). VERSACE, Richard, William [US/US]; 69 Townsend Road, Wanaque, NJ 07465 (US). SHARMA, Sushil, Kumar [US/US]; 9 Bakley Terrace, West Orange, NJ 07052 (US).

- (74) Agent: BECKER, Konrad; Novartis AG, Corporate Intellectual Property, Patent & Trademark Department, CH-4002 Basel (CH).
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(54) Title: DEACETYLASE INHIBITORS

(57) Abstract: The present invention provides hydroxamate compounds which are deacetylase inhibitors. The compounds are suitable for pharmaceutical compositions having anti-proliferative properties.

DEACETYLASE INHIBITORS

The present invention relates to hydroxamate compounds which are inhibitors of histone deacetylase. The inventive compounds are useful as pharmaceuticals for the treatment of proliferative diseases.

Background

Reversible acetylation of histones is a major regulator of gene expression that acts by altering accessibility of transcription factors to DNA. In normal cells, histone deacetylase (HDA) and histone acetyltrasferase together control the level of acetylation of histones to maintain a balance. Inhibition of HDA results in the accumulation of hyperacetylated histones, which results in a variety of cellular responses.

Inhibitors of HDA have been studied for their therapeutic effects on cancer cells. For example, butyric acid and its derivatives, including sodium phenylbutyrate, have been reported to induce apoptosis *in vitro* in human colon carcinoma, leukemia and retinoblastoma cell lines. However, butyric acid and its derivatives are not useful pharmacological agents because they tend to be metabolized rapidly and have a very short half-life *in vivo*. Other inhibitors of HDA that have been widely studied for their anti-cancer activities are trichostatin A and trapoxin. Trichostatin A is an antifungal and antibiotic and is a reversible inhibitor of mammalian HDA. Trapoxin is a cyclic tetrapeptide, which is an irreversible inhibitor of mammalian HDA. Although trichostatin and trapoxin have been studied for their anti-cancer activities, the *in vivo* instability of the compounds makes them less suitable as anti-cancer drugs. There remains a need for an active compound that is suitable for treating tumors, including cancerous tumors, that is highly efficacious and stable.

Summary

The present invention provides efficacious deacetylase inhibitor compounds that are useful as pharmaceutical agents having the formula I

HO
$$R_1$$
 R_2 R_3 R_4 R_5 R_5 R_5

wherein

- R₁ is H, halo, or a straight chain C₁-C₆ alkyl (especially methyl, ethyl or *n*-propyl, which methyl, ethyl and n-propyl substituents are unsubstituted or substituted by one or more substituents described below for alkyl substituents);
- R_2 is selected from H, C_1 - C_{10} alkyl, (e.g. methyl, ethyl or - CH_2CH_2 -OH), C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, C_4 C_9 heterocycloalkylalkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), aryl, heteroaryl, arylalkyl (e.g. benzyl), heteroarylalkyl (e.g. pyridylmethyl), - $(CH_2)_nC(O)R_6$, - $(CH_2)_nOC(O)R_6$, amino acyl, HON-C(O)-CH=C(R_1)-aryl-alkyl- and - $(CH_2)_nR_7$;
- R₃ and R₄ are the same or different and independently H, C₁-C₆ alkyl, acyl or acylamino, or R₃ and R₄ together with the carbon to which they are bound represent C=O, C=S, or C=NR₈, or R₂ together with the nitrogen to which it is bound and R₃ together with the carbon to which it is bound can form a C₄ C₉ heterocycloalkyl, a heteroaryl, a polyheteroaryl, a non-aromatic polyheterocycle, or a mixed aryl and non-aryl polyheterocycle ring;
- R₅ is selected from H, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, acyl, aryl, heteroaryl, arylalkyl (e.g. benzyl), heteroarylalkyl (e.g. pyridylmethyl), aromatic polycycles, non-aromatic polycycles, mixed aryl and non-aryl polycycles, polyheteroaryl, non-aromatic polyheterocycles, and mixed aryl and non-aryl polyheterocycles;
- n_1 , n_2 and n_3 are the same or different and independently selected from 0-6, when n_1 is 1-6, each carbon atom can be optionally and independently substituted with R_3 and/or R_4 ;
- X and Y are the same or different and independently selected from H, halo, C₁-C₄ alkyl, such as CH₃ and CF₃, NO₂, C(O)R₁, OR₉, SR₉, CN, and NR₁₀R₁₁;

- R_6 is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), aryl, heteroaryl, arylalkyl (e.g., benzyl, 2-phenylethenyl), heteroarylalkyl (e.g., pyridylmethyl), OR_{12} , and OR_{13} R₁₄;
- R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{18}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_6$;
- R_8 is selected from H, OR_{15} , $NR_{13}R_{14}$, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl);
- R_9 is selected from $C_1 C_4$ alkyl, for example, CH_3 and CF_3 , C(O)-alkyl, for example $C(O)CH_3$, and $C(O)CF_3$;
- R₁₀ and R₁₁ are the same or different and independently selected from H, C₁-C₄ alkyl, and -C(O)-alkyl;
- R₁₂ is selected from H, C₁-C₈ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, C₄ C₉ heterocycloalkylalkyl, aryl, mixed aryl and non-aryl polycycle, heteroaryl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl);
- R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), heteroarylalkyl (e.g., pyridylmethyl), amino acyl, or R_{13} and R_{14} together with the nitrogen to which they are bound are C_4 C_9 heterocycloalkyl, heteroaryl, polyheteroaryl, non-aromatic polyheterocycle or mixed aryl and non-aryl polyheterocycle;
- R_{15} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{16} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, polyheteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{17} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, aromatic polycycles, heteroaryl, arylalkyl, heteroarylalkyl, polyheteroaryl and $NR_{13}R_{14}$;
- m is an integer selected from 0 to 6; and
- Z is selected from O, NR₁₃, S and S(O),
- or a pharmaceutically acceptable salt thereof.

The compounds of the present invention are suitable as active agents in pharmaceutical compositions that are efficacious particularly for treating cellular proliferative ailments. The pharmaceutical composition has a pharmaceutically effective amount of the

present active agent along with other pharmaceutically acceptable exipients, carriers, fillers, diluents and the like. The term pharmaceutically effective amount as used herein indicates an amount necessary to administer to a host to achieve a therapeutic result, especially an anti-tumor effect, e.g., inhibition of proliferation of malignant cancer cells, benign tumor cells or other proliferative cells.

Detailed Description

The present invention provides hydroxamate compounds, e.g., hydroxamic acids, that are inhibitors of deacetylases, preferably inhibitors of histone deacetylases. The hydroxamate compounds are highly suitable for treating tumors, including cancerous tumors. The hydroxamate compounds of the present invention have the following structure I

HO
$$R_1$$
 R_2 R_3 R_4 R_5 R_5 R_5

wherein

R₁ is H, halo, or a straight chain C₁-C₈ alkyl (especially methyl, ethyl or *n*-propyl, which methyl, ethyl and n-propyl substituents are unsubstituted or substituted by one or more substituents described below for alkyl substituents);

R₂ is selected from H, C₁-C₁₀ alkyl, (preferably C₁-C₆ alkyl, e.g. methyl, ethyl or -CH₂CH₂-OH), C₄ - C₉ cycloalkyl, C₄ - C₉ heterocycloalkyl, C₄ - C₉ heterocycloalkylalkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), aryl, heteroaryl, arylalkyl (e.g. benzyl), heteroarylalkyl (e.g. pyridylmethyl), -(CH₂)_nC(O)R₆, -(CH₂)_nOC(O)R₆, amino acyl, HON-C(O)-CH=C(R₁)-aryl-alkyl- and -(CH₂)_nR₇;

R₃ and R₄ are the same or different and independently H, C₁-C₆ alkyl, acyl or acylamino, or R₃ and R₄ together with the carbon to which they are bound represent C=O, C=S, or C=NR₈, or R₂ together with the nitrogen to which it is bound and R₃ together with the carbon to which it is bound can form a C₄ - C₉ heterocycloalkyl, a

- heteroaryl, a polyheteroaryl, a non-aromatic polyheterocycle, or a mixed aryl and non-aryl polyheterocycle ring;
- R₅ is selected from H, C₁-C₈ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, acyl, aryl, heteroaryl, arylalkyl (e.g. benzyl), heteroarylalkyl (e.g. pyridylmethyl), aromatic polycycles, non-aromatic polycycles, mixed aryl and non-aryl polycycles, polyheteroaryl, non-aromatic polyheterocycles, and mixed aryl and non-aryl polyheterocycles;
- n_1 , n_2 and n_3 are the same or different and independently selected from 0-6, when n_1 is 1-6, each carbon atom can be optionally and independently substituted with R_3 and/or R_4 ;
- X and Y are the same or different and independently selected from H, halo, C₁-C₄ alkyl, such as CH₃ and CF₃, NO₂, C(O)R₁, OR₉, SR₉, CN, and NR₁₀R₁₁;
- R₆ is selected from H, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), aryl, heteroaryl, arylalkyl (e.g., benzyl, 2-phenylethenyl), heteroarylalkyl (e.g., pyridylmethyl), OR₁₂, and NR₁₃R₁₄;
- R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{16}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_{6}$;
- R_8 is selected from H, OR_{15} , $NR_{13}R_{14}$, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl);
- R_9 is selected from $C_1 C_4$ alkyl, for example, CH_3 and CF_3 , C(O)-alkyl, for example $C(O)CH_3$, and $C(O)CF_3$;
- R₁₀ and R₁₁ are the same or different and independently selected from H, C₁-C₄ alkyl, and -C(O)-alkyl;
- R_{12} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, C_4 C_9 heterocycloalkylalkyl, aryl, mixed aryl and non-aryl polycycle, heteroaryl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl);
- R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), heteroarylalkyl (e.g., pyridylmethyl), amino acyl, or R_{13} and R_{14} together with the nitrogen to which they are bound are C_4 C_9 heterocycloalkyl, heteroaryl, polyheteroaryl, non-aromatic polyheterocycle or mixed aryl and non-aryl polyheterocycle;
- R_{15} is selected from H, C_1 - C_8 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;

 R_{16} is selected from C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, polyheteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;

R₁₇ is selected from C₁-C₆ alkyl, C₄ – C₉ cycloalkyl, C₄ – C₉ heterocycloalkyl, aryl, aromatic polycycles, heteroaryl, arylalkyl, heteroarylalkyl, polyheteroaryl and NR₁₃R₁₄;

m is an integer selected from 0 to 6; and Z is selected from O, NR₁₃, S and S(O), or a pharmaceutically acceptable salt thereof.

As appropriate, unsubstituted means that there is no substituent or that the only substituents are hydrogen.

Halo substituents are selected from fluoro, chloro, bromo and iodo, preferably fluoro or chloro.

Alkyl substituents include straight and branched C₁-C₆alkyl, unless otherwise noted. Examples of suitable straight and branched C₁-C₆alkyl substituents include methyl, ethyl, n-propyl, 2-propyl, n-butyl, sec-butyl, t-butyl, and the like. Unless otherwise noted, the alkyl substituents include both unsubstituted alkyl groups and alkyl groups that are substituted by one or more suitable substituents, including unsaturation (i.e. there are one or more double or triple C-C bonds), acyl, cycloalkyl, halo, oxyalkyl, alkylamino, aminoalkyl, acylamino and OR₁₅, for example, alkoxy. Preferred substituents for alkyl groups include halo, hydroxy, alkoxy, oxyalkyl, alkylamino, and aminoalkyl.

Cycloalkyl substituents include C_3 - C_9 cycloalkyl groups, such as cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl and the like, unless otherwise specified. Unless otherwise noted, cycloalkyl substituents include both unsubstituted cycloalkyl groups and cycloalkyl groups that are substituted by one or more suitable substituents, including C_1 - C_9 alkyl, halo, hydroxy, aminoalkyl, oxyalkyl, alkylamino, and OR_{15} , such as alkoxy. Preferred substituents for cycloalkyl groups include halo, hydroxy, alkoxy, oxyalkyl, alkylamino and aminoalkyl.

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The above discussion of alkyl and cycloalkyl substituents also applies to the alkyl portions of other substituents, such as without limitation, alkoxy, alkyl amines, alkyl ketones, arylalkyl, heteroarylalkyl, alkylsulfonyl and alkyl ester substituents and the like.

Heterocycloalkyl substituents include 3 to 9 membered aliphatic rings, such as 4 to 7 membered aliphatic rings, containing from one to three heteroatoms selected from nitrogen, sulfur and oxygen. Examples of suitable heterocycloalkyl substituents include pyrrolidyl, tetrahydrofuryl, tetrahydrothiofuranyl, piperidyl, piperazyl, tetrahydropyranyl, morphilino, 1,3-diazapane, 1,4-diazapane, 1,4-oxazepane, and 1,4-oxathiapane. Unless otherwise noted, the rings are unsubstituted or substuted on the carbon atoms by one or more suitable substituents, including C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl), halo, amino, alkyl amino and OR_{15} , for example alkoxy. Unless otherwise noted, nitrogen heteroatoms are unsubstituted or substituted by H, C_1 - C_4 alkyl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl), acyl, aminoacyl, alkylsulfonyl, and arylsulfonyl.

Cycloalkylalkyl substituents include compounds of the formula –(CH₂)_{n5}-cycloalkyl wherein n5 is a number from 1-6. Suitable cycloalkylalkyl substituents include cyclopentylmethyl-, cyclopentylethyl, cyclohexylmethyl and the like. Such substituents are unsubstituted or substituted in the alkyl portion or in the cycloalkyl portion by a suitable substituent, including those listed above for alkyl and cycloalkyl.

Aryl substituents include unsubstituted phenyl and phenyl substituted by one or more suitable substituents, including C₁-C₆ alkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), O(CO)alkyl, oxyalkyl, halo, nitro, amino, alkylamino, aminoalkyl, alkyl ketones, nitrile, carboxyalkyl, alkylsulfonyl, aminosulfonyl, arylsulfonyl, and OR₁₅, such as alkoxy. Preferred substituents include including C₁-C₆ alkyl, cycloalkyl (e.g., cyclopropylmethyl), alkoxy, oxyalkyl, halo, nitro, amino, alkylamino, aminoalkyl, alkyl ketones, nitrile, carboxyalkyl, alkylsulfonyl, arylsulfonyl, and aminosulfonyl. Examples of suitable aryl groups include C₁-C₄alkylphenyl, C₁-C₄alkoxyphenyl, trifluoromethylphenyl, methoxyphenyl, hydroxyethylphenyl, dimethylaminophenyl, aminopropylphenyl, carbethoxyphenyl, methanesulfonylphenyl and tolylsulfonylphenyl.

Aromatic polycycles include naphthyl, and naphthyl substituted by one or more suitable substituents, including C_1 - C_8 alkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), oxyalkyl, halo, nitro, amino, alkylamino, aminoalkyl, alkyl ketones, nitrile, carboxyalkyl, alkylsulfonyl, arylsulfonyl, aminosulfonyl and OR_{15} , such as alkoxy.

Heteroaryl substituents include compounds with a 5 to 7 member aromatic ring containing one or more heteroatoms, for example from 1 to 4 heteroatoms, selected from N, O and S. Typical heteroaryl substituents include furyl, thienyl, pyrrole, pyrazole, triazole, thiazole, oxazole, pyridine, pyrimidine, isoxazolyl, pyrazine and the like. Unless otherwise noted, heteroaryl substituents are unsubstituted or substituted on a carbon atom by one or more suitable substituents, including alkyl, the alkyl substituents identified above, and another heteroaryl substituent. Nitrogen atoms are unsubstituted or substituted, for example by R_{13} ; especially useful N substituents include H, $C_1 - C_4$ alkyl, acyl, aminoacyl, and sulfonyl.

Arylalkyl substituents include groups of the formula $-(CH_2)_{n5}$ -aryl, $-(CH_2)_{n5-1}$ - $(CHaryl)-(CH_2)_{n5}$ -aryl or $-(CH_2)_{n5-1}$ CH(aryl)(aryl) wherein aryl and n5 are as defined above. Such arylalkyl substituents include benzyl, 2-phenylethyl, 1-phenylethyl, tolyl-3-propyl, 2-phenylpropyl, diphenylmethyl, 2-diphenylethyl, 5,5-dimethyl-3-phenylpentyl and the like. Arylalkyl substituents are unsubstituted or substituted in the alkyl moiety or the aryl moiety or both as described above for alkyl and aryl substituents.

Heteroarylalkyl substituents include groups of the formula –(CH₂)_{n5}-heteroaryl wherein heteroaryl and n5 are as defined above and the bridging group is linked to a carbon or a nitrogen of the heteroaryl portion, such as 2-, 3- or 4-pyridylmethyl, imidazolylmethyl, quinolylethyl, and pyrrolylbutyl. Heteroaryl substituents are unsubstituted or substituted as discussed above for heteroaryl and alkyl substituents.

Amino acyl substituents include groups of the formula $-C(O)-(CH_2)_n-C(H)(NR_{13}R_{14})-(CH_2)_n-R_5$ wherein n, R_{13} , R_{14} and R_5 are described above. Suitable aminoacyl substituents include natural and non-natural amino acids such as glycinyl, D-tryptophanyl, L-lysinyl, D- or L-homoserinyl, 4-aminobutryic acyl, \pm -3-amin-4-hexenoyl.

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Non-aromatic polycycle substituents include bicyclic and tricyclic fused ring systems where each ring can be 4-9 membered and each ring can contain zero, 1 or more double and/or triple bonds. Suitable examples of non-aromatic polycycles include decalin, octahydroindene, perhydrobenzocycloheptene, perhydrobenzo-[f]-azulene. Such substituents are unsubstituted or substituted as described above for cycloalkyl groups.

Mixed aryl and non-aryl polycycle substituents include bicyclic and tricyclic fused ring systems where each ring can be 4 – 9 membered and at least one ring is aromatic. Suitable examples of mixed aryl and non-aryl polycycles include methylenedioxyphenyl, bismethylenedioxyphenyl, 1,2,3,4-tetrahydronaphthalene, dibenzosuberane, dihdydroanthracene, 9H-fluorene. Such substituents are unsubstituted or substituted by nitro or as described above for cycloalkyl groups.

Polyheteroaryl substituents include bicyclic and tricyclic fused ring systems where each ring can independently be 5 or 6 membered and contain one or more heteroatom, for example, 1, 2, 3, or 4 heteroatoms, chosen from O, N or S such that the fused ring system is aromatic. Suitable examples of polyheteroaryl ring systems include quinoline, isoquinoline, pyridopyrazine, pyrrolopyridine, furopyridine, indole, benzofuran, benzothiofuran, benzindole, benzoxazole, pyrroloquinoline, and the like. Unless otherwise noted, polyheteroaryl substituents are unsubstituted or substituted on a carbon atom by one or more suitable substituents, including alkyl, the alkyl substituents identified above and a substituent of the formula -O-($CH_2CH=CH(CH_3)(CH_2)$)₁₋₃H. Nitrogen atoms are unsubstituted or substituted, for example by R_{13} ; especially useful N substituents include H, $C_1 - C_4$ alkyl, acyl, aminoacyl, and sulfonyl.

Non-aromatic polyheterocyclic substituents include bicyclic and tricyclic fused ring systems where each ring can be 4 – 9 membered, contain one or more heteroatom, for example, 1, 2, 3, or 4 heteroatoms, chosen from O, N or S and contain zero or one or more C-C double or triple bonds. Suitable examples of non-aromatic polyheterocycles include hexitol, cis-perhydro-cyclohepta[b]pyridinyl, decahydro-benzo[f][1,4]oxazepinyl, 2,8-dioxabicyclo[3.3.0]octane, hexahydro-thieno[3,2-b]thiophene, perhydropyrrolo[3,2-b]pyrrole, perhydronaphthyridine, perhydro-1H-dicyclopenta[b,e]pyran. Unless otherwise noted, non-aromatic polyheterocyclic substituents are unsubstituted or substituents identified above.

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Nitrogen atoms are unsubstituted or substituted, for example, by R_{13} ; especially useful N substituents include H, $C_1 - C_4$ alkyl, acyl, aminoacyl, and sulfonyl.

Mixed aryl and non-aryl polyheterocycles substituents include bicyclic and tricyclic fused ring systems where each ring can be 4 – 9 membered, contain one or more heteroatom chosen from O, N or S, and at least one of the rings must be aromatic. Suitable examples of mixed aryl and non-aryl polyheterocycles include 2,3-dihydroindole, 1,2,3,4-tetrahydroquinoline, 5,11-dihydro-10H-dibenz[b,e][1,4]diazepine, 5H-dibenzo[b,e][1,4]diazepine, 1,2-dihydropyrrolo[3,4-b][1,5]benzodiazepine, 1,5-dihydropyrido[2,3-b][1,4]diazepin-4-one, 1,2,3,4,6,11-hexahydro-benzo[b]pyrido[2,3-e][1,4]diazepin-5-one. Unless otherwise noted, mixed aryl and non-aryl polyheterocyclic substituents are unsubstituted or substituted on a carbon atom by one or more suitable substituents, including, -N-OH, =N-OH, alkyl and the alkyl substituents identified above. Nitrogen atoms are unsubstituted or substituted, for example, by R₁₃; especially useful N substituents include H, C₁ – C₄ alkyl, acyl, aminoacyl, and sulfonyl.

Amino substituents include primary, secondary and tertiary amines and in salt form, quaternary amines. Examples of amino substituents include mono- and di-alkylamino, mono- and di-aryl amino, mono- and di-arylalkylamino, alkyl-arylalkylamino and the like.

Sulfonyl substituents include alkylsulfonyl and arylsulfonyl, for example methane sulfonyl, benzene sulfonyl, tosyl and the like.

Acyl substituents include groups of the formula –C(O)-W, –OC(O)-W, –C(O)-O-W and –C(O)NR₁₃R₁₄, where W is R₁₆, H or cycloalkylalkyl.

Acylamino substituents include groups of the formula $-N(R_{12})C(O)-W$, $-N(R_{12})C(O)-W$, and $-N(R_{12})C(O)-NHOH$ and R_{12} and W are as defined above.

The R₂ substituent HON-C(O)-CH=C(R₁)-aryl-alkyl- is a group of the formula

wherein n₄ is 0-3 and X and Y are as defined above.

Preferences for each of the substituents include the following:

R₁ is H, halo, or a straight chain C₁-C₄ alkyl;

 R_2 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, amino acyl, and -(CH₂)_nR₇;

R₃ and R₄ are the same or different and independently selected from H, and C₁-C₆ alkyl, or R₃ and R₄ together with the carbon to which they are bound represent C=O, C=S, or C=NR₈;

R₅ is selected from H, C₁-C₆ alkyl, C₄ – C₉ cycloalkyl, C₄ – C₉ heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, an aromatic polycycle, a non-aromatic polycycle, a mixed aryl and non-aryl polycycle, polyheteroaryl, a non-aromatic polyheterocycle, and a mixed aryl and non-aryl polyheterocycle;

 n_1 , n_2 and n_3 are the same or different and independently selected from 0-6, when n_1 is 1-6, each carbon atom is unsubstituted or independently substituted with R_3 and/or R_4 ;

X and Y are the same or different and independently selected from H, halo, C_1 - C_4 alkyl, CF_3 , NO_2 , $C(O)R_1$, OR_9 , SR_9 , CN, and $NR_{10}R_{11}$;

 R_6 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, OR_{12} , and $NR_{13}R_{14}$;

 R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{16}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_{6}$;

 R_8 is selected from H, OR_{15} , $NR_{13}R_{14}$, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;

 R_{θ} is selected from $C_1 - C_4$ alkyl and C(O)-alkyl;

 R_{10} and R_{11} are the same or different and independently selected from H, C_1 - C_4 alkyl, and -C(O)-alkyl;

 R_{12} is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;

 R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and amino acyl;

 R_{15} is selected from H, C_1 - C_8 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;

 R_{16} is selected from C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;

 R_{17} is selected from C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $NR_{13}R_{14}$;

m is an integer selected from 0 to 6; and

Z is selected from O, NR₁₃, S, S(O).

Useful compounds of the formula I include those wherein each of R_1 , X, Y, R_3 , and R_4 is H, including those wherein one of n_2 and n_3 is zero and the other is 1, especially those wherein R_2 is H or -CH₂-CH₂-OH.

One suitable genus of hydroxamate compounds are those of formula la

HO
$$R_2$$
 (la)

wherein

n₄ is 0-3,

R₂ is selected from H, C₁-C₆ alkyl, C₄ – C₉ cycloalkyl, C₄ – C₉ heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, amino acyl and -(CH₂)_nR₇;

R₅' is heteroaryl, heteroarylalkyl (e.g., pyridylmethyl), aromatic polycycles, non-aromatic polycycles, mixed aryl and non-aryl polycycles, polyheteroaryl, or mixed aryl and non-aryl polyheterocycles,

or a pharmaceutically acceptable salt thereof.

Another suitable genus of hydroxamate compounds are those of formula la

HO
$$R_2$$
 (Ia)

wherein

n₄ is 0-3,

 R_2 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, amino acyl and -(CH₂)_nR₇;

R₅' is aryl, arylalkyl, aromatic polycycles, non-aromatic polycycles, and mixed aryl and non-aryl polycycles; especially aryl, such as p-fluorophenyl, p-chlorophenyl, p-O-C₁-C₄-alkylphenyl, such as p-methoxyphenyl, and p-C₁-C₄-alkylphenyl; and arylalkyl, such as benzyl, ortho, meta or para-fluorobenzyl, ortho, meta or para-chlorobenzyl, ortho, meta or para-mono, di or tri-O-C₁-C₄-alkylbenzyl, such as ortho, meta or para-methoxybenzyl, m,p-dlethoxybenzyl, o,m,p-triimethoxybenzyl, and ortho, meta or para-mono, di or tri C₁-C₄-alkylphenyl, such as p-methyl, m,m-dlethylphenyl, or a pharmaceutically acceptable salt thereof.

Another interesting genus are the compounds of formula lb

$$\begin{array}{c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

wherein

 R_2 ' is selected from H, C_1 - C_6 alkyl, C_4 - C_6 cycloalkyl, cycloalkylalkyl (e.g., cyclopropylmethyl), -(CH_2)₂₋₄ OR_{21} where R_{21} is H, methyl, ethyl, propyl, and *i*-propyl, and

 R_5 " is unsubstituted 1*H*-indol-3-yl, benzofuran-3-yl or quinolin-3-yl, or substituted 1*H*-indol-3-yl, such as 5-fluoro-1*H*-indol-3-yl or 5-methoxy-1*H*-indol-3-yl, benzofuran-3-yl or quinolin-3-yl,

or a pharmaceutically acceptable salt thereof.

Another interesting genus of hydroxamate compounds are the compounds of formula Ic

HO
$$R_1$$
 R_{18} R_{18} R_{18} R_{19} R

wherein

the ring containing Z_1 is aromatic or non-aromatic, which non-aromatic rings are saturated or unsaturated,

Z₁ is O, S or N-R₂₀,

R₁₈ is H, halo, C₁-C₆alkyl (methyl, ethyl, t-butyl), C₃-C₇cycloalkyl, aryl, for example unsubstituted phenyl or phenyl substituted by 4-OCH₃ or 4-CF₃, or heteroaryl, such as 2-furanyl, 2-thiophenyl or 2-, 3- or 4-pyridyl;

 R_{20} is H, C_1 - C_6 alkyl, C_1 - C_6 alkyl- C_3 - C_9 cycloalkyl (e.g., cyclopropylmethyl), aryl, heteroaryl, arylalkyl (e.g., benzyl), heteroarylalkyl (e.g., pyridylmethyl), acyl (acetyl, propionyl, benzoyl) or sulfonyl (methanesulfonyl, ethanesulfonyl, benzenesulfonyl, toluenesulfonyl);

A₁ is 1, 2 or 3 substituents which are independently H, C₁-C-₆alkyl, -OR₁₉, halo, alkylamino, aminoalkyl, halo, or heteroarylalkyl (e.g., pyridylmethyl),

 R_{19} is selected from H, C_1 - C_6 alkyl, C_4 - C_9 cycloalkyl, C_4 - C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), heteroarylalkyl (e.g., pyridylmethyl) and

-(CH₂CH=CH(CH₃)(CH₂))₁₋₃H;

 R_2 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, amino acyl and -(CH₂)_nR₇;

v is 0, 1 or 2,

p is 0-3, and

q is 1-5 and r is 0 or

q is 0 and r is 1-5,

or a pharmaceutically acceptable salt thereof. The other variable substituents are as defined above.

Especially useful compounds of formula Ic are those wherein R_2 is H, or $-(CH_2)_pCH_2OH$, wherein p is 1-3, especially those wherein R_1 is H; such as those wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3, especially those wherein Z_1 is N-R₂₀. Among these compounds R_2 is preferably H or $-CH_2-CH_2-OH$ and the sum of q and r is preferably 1.

Another interesting genus of hydroxamate compounds are the compounds of formula Id

HO
$$R_1$$
 R_{1B} R

wherein

Z₁ is O, S or N-R₂₀,

 R_{18} is H, halo, C_1 - C_6 alkyl (methyl, ethyl, t-butyl), C_3 - C_7 cycloalkyl, aryl, for example, unsubstituted phenyl or phenyl substituted by 4-OCH₃ or 4-CF₃, or heteroaryl,

 R_{20} is H, C_1 - C_6 alkyl, C_1 - C_6 alkyl- C_3 - C_9 cycloalkyl (e.g., cyclopropylmethyl), aryl, heteroaryl, arylalkyl (e.g., benzyl), heteroarylalkyl (e.g., pyridylmethyl), acyl (acetyl, propionyl, benzoyl) or sulfonyl (methanesulfonyl, ethanesulfonyl, benzenesulfonyl, toluenesulfonyl); A₁ is 1, 2 or 3 substituents which are independently H, C_1 - C_6 alkyl, - OR_{19} , or halo, R_{19} is selected from H, C_1 - C_6 alkyl, C_4 - C_9 cycloalkyl, C_4 - C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl (e.g., benzyl), and heteroarylalkyl (e.g., pyridylmethyl); p is 0-3, and

p 13 0-0, and

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q is 1-5 and r is 0 or

q is 0 and r is 1-5,

or a pharmaceutically acceptable salt thereof. The other variable substituents are as defined above.

Especially useful compounds of formula Id are those wherein R_2 is H, or - $(CH_2)_pCH_2OH$, wherein p is 1-3, especially those wherein R_1 is H; such as those wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3. Among these compounds R_2 is preferably H or - CH_2 - CH_2 -OH and the sum of q and r is preferably 1.

The present invention further relates to compounds of the formula le

HO
$$R_1$$
 R_2 R_3 R_4 N - R_{20} (le)

or a pharmaceutically acceptable salt thereof. The variable substituents are as defined above.

Especially useful compounds of formula le are those wherein R₁₈ is H, fluoro, chloro, bromo, a C₁-C₄alkyl group, a substituted C₁-C₄alkyl group, a C₃-C₇cycloalkyl group, unsubstituted phenyl, phenyl substituted in the para position, or a heteroaryl (e.g., pyridyl) ring.

Another group of useful compounds of formula le are those wherein R_2 is H, or - $(CH_2)_pCH_2OH$, wherein p is 1-3, especially those wherein R_1 is H; such as those wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3. Among these compounds R_2 is preferably H or $-CH_2-CH_2-OH$ and the sum of q and r is preferably 1.

Another group of useful compounds of formula le are those wherein R_{18} is H, methyl, ethyl, t-butyl, trifluoromethyl, cyclohexyl, phenyl, 4-methoxyphenyl, 4-trifluoromethylphenyl, 2-furanyl, 2-thiophenyl, or 2-, 3- or 4-pyridyl wherein the 2-furanyl, 2-thiophenyl and 2-, 3- or 4-pyridyl substituents are unsubstituted or substituted as described above for heteroaryl rings; R_2 is H, or -(CH₂)_pCH₂OH, wherein p is 1-3; especially those wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3. Among these compounds R_2 is preferably H or -CH₂-CH₂-OH and the sum of q and r is preferably 1.

Those compounds of formula le wherein R₂₀ is H or C₁-C₆alkyl, especially H, are important members of each of the subgenuses of compounds of formula le described above.

N-hydroxy-3-[4-[[(2-hydroxyethyl)[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, N-hydroxy-3-[4-[[[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide and N-hydroxy-3-[4-[[[2-(2-methyl-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide, or a pharmaceutically acceptable salt thereof, are important compounds of formula le.

The present invention further relates to the compounds of the formula If

HO N
$$\mathbb{R}_1$$
 \mathbb{R}_2 \mathbb{R}_3 \mathbb{R}_4 (Iff)

or a pharmaceutically acceptable salt thereof. The variable substituents are as defined above.

Useful compounds of formula If are those wherein R_2 is H, or -(CH_2) $_pCH_2OH$, wherein p is 1-3, especially those wherein R_1 is H; such as those wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3. Among these compounds R_2 is preferably H or - CH_2 - CH_2 -OH and the sum of q and r is preferably 1.

N-hydroxy-3-[4-[[[2-(benzofur-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide,or a pharmaceutically acceptable salt thereof, is an important compound of formula If.

The compounds described above are often used in the form of a pharmaceutically acceptable salt. Pharmaceutically acceptable salts include, when appropriate, pharmaceutically acceptable base addition salts and acid addition salts, for example, metal salts, such as alkali and alkaline earth metal salts, ammonium salts, organic amine addition salts, and amino acid addition salts, and sulfonate salts. Acid addition salts include inorganic acid addition salts such as hydrochloride, sulfate and phosphate, and organic acid addition salts such as alkyl sulfonate, arylsulfonate, acetate, maleate, fumarate, tartrate, citrate and lactate. Examples of metal salts are alkali metal salts, such as lithium salt, sodium salt and potassium salt, alkaline earth metal salts such as magnesium salt and calcium salt, aluminum salt, and zinc salt. Examples of ammonium salts are ammonium salt and tetramethylammonium salt. Examples of organic amine addition salts are salts with morpholine and piperidine. Examples of amino acid addition salts are salts with glycine, phenylalanine, glutamic acid and lysine. Sulfonate salts include mesylate, tosylate and benzene sulfonic acid salts.

As is evident to those skilled in the art, the many of the deacetylase inhibitor compounds of the present invention contain asymmetric carbon atoms. It should be understood, therefore, that the individual stereoisomers are contemplated as being included within the scope of this invention.

The hydroxamate compounds of the present invention can be produced by known organic synthesis methods. For example, the hydroxamate compounds can be produced

by reacting methyl 4-formyl cinnamate with tryptamine and then converting the reactant to the hydroxamate compounds. As an example, methyl 4-formyl cinnamate 2, is prepared by acid catalyzed esterification of 4-formylcinnamic acid 3 (Bull. Chem. Soc. Jpn. 1995; 68:2355-2362). An alternate preparation of methyl 4-formyl cinnamate 2 is by a Pd-catalyzed coupling of methyl acrylate 4 with 4-bromobenzaldehyde 5.

Additional starting materials can be prepared from 4-carboxybenzaldehyde 6, and an exemplary method is illustrated for the preparation of aldehyde 9, shown below. The carboxylic acid in 4-carboxybenzaldehyde 6 can be protected as a silyl ester (e.g., the t-butyldimethylsilyl ester) by treatment with a silyl chloride (e.g., t-butyldimethylsilyl chloride) and a base (e.g. triethylamine) in an appropriate solvent (e.g., dichloromethane). The resulting silyl ester 7 can undergo an olefination reaction (e.g., a Horner-Emmons olefination) with a phosphonate ester (e.g., triethyl 2-phosphonopropionate) in the presence of a base (e.g., sodium hydride) in an appropriate solvent (e.g., tetrahydrofuran (THF)). Treatment of the resulting diester with acid (e.g., aqueous hydrochloric acid) results in the hydrolysis of the silyl ester providing acid 8. Selective reduction of the carboxylic acid of 8 using, for example, borane-dimethylsuflide complex in a solvent (e.g., THF) provides an intermediate alcohol. This intermediate alcohol could be oxidized to aldehyde 9 by a number of known methods, including, but not limited to, Swern oxidation, Dess-Martin periodinane oxidation, Moffatt oxidation and the like.

The aldehyde starting materials 2 or 9 can be reductively aminated to provide secondary or tertiary amines. This is illustrated by the reaction of methyl 4-formyl cinnamate 2 with tryptamine 10 using sodium triacetoxyborohydride (NaBH(OAc)₃) as the reducing agent in dichloroethane (DCE) as solvent to provide amine 11. Other reducing agents can be used, e.g., sodium borohydride (NaBH₄) and sodium cyanoborohydride (NaBH₃CN), in other solvents or solvent mixtures in the presence or absence of acid catalysts (e.g., acetic acid and trifluoroacetic acid). Amine 11 can be converted directly to hydroxamic acid 12 by treatment with 50% aqueous hydroxylamine in a suitable solvent (e.g., THF in the presence of a base, e.g., NaOH). Other methods of hydroxamate formation are known and include reaction of an ester with hydroxylamine hydrochloride and a base (e.g., sodium hydroxide or sodium methoxide) in a suitable solvent or solvent mixture (e.g., methanol, ethanol or methanol/THF).

Aldehyde 2 can be reductively aminated with a variety of amines, exemplified by, but not limited to, those illustrated in Table 1. The resulting esters can be converted to target hydroxamates by the methods listed.

Table 1

Amine	Reducing	Hydroxamate	R
	Conditions	Conditions	·

	T		·
NH ₂	NaBH(OAc)₃	2 M HONH ₂ in	CH₂
N N	HOAc, DCE	MeOH	\ \mathrea{\text{N}}
HN NH ₂	65	cs	HN CH ₂
U _N NH₂	st .	a a	CIN CH2
NH ₂	и	cf	CH ₂
NH ₂	4	ıı	F CH₂
MeO NH ₂		tt	MeO CH ₂
SO ₂ HN-NH ₂	и	cr.	SO ₂ HN— CH ₂
NH ₂	а	ď	NN CH₂ Me
N√NH₂	æ	u	SN_CH ₂
Ph(CH₂)₃NH₂	NaBH₃CN/MeOH/ HOAc		Ph(CH₂)₃

An alternate synthesis of the compounds of this invention starts by reductive amination of 4-formyl cinnamic acid 3, illustrated below with 3-phenylpropylamine 13, using, for example, NaBH₃CN as the reducing agent in MeOH and HOAc as a catalyst. The basic nitrogen of the resulting amino acid 14 can be protected, for example, as *t*-butoxycarbamate (BOC) by reaction with di-*t*-butyldicarbonate to give 15.

The carboxylic acid can be coupled with a protected hydroxylamine (e.g., *O*-trityl hydroxylamine) using a dehydrating agent (e.g., 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDCI)) and a catalyst (e.g., 1-hydroxybenzotriazole hydrate (HOBT)) in a suitable solvent (e.g., DMF) to produce **16**. Treatment of **16** with a strong acid (e.g., trifluoroacetic acid (TFA)) provides a hydroxamic acid **17** of the present invention. Additional examples of compounds that can be prepared by this method are:

Tertiary amine compounds can be prepared by a number of methods. Reductive amination of **30** with nicotinaldehyde **32** using NaBH₃CN as the reducing agent in dichloroethane and HOAc as a catalyst provides ester **34**. Other reducing agents can be used (e.g., NaBH₄ and NaBH(OAc)₃) in other solvents or solvent mixtures in the presence or absence of acid catalysts (e.g., acetic acid, trifluoroacetic acid and the like). Reaction of ester **34** with HONH₂•HCI, NaOH in MeOH provides hydroxamate **36**.

Tertiary amine compounds prepared by this methodology are exemplified, but not limited to, those listed in Table 2.

Table 2

	Reducing Conditions	Hydroxamate
		Conditions
CH ₂	NaBH(OAc)₃ HOAc,	HONH ₂ •HCI/NaOMe/
N.	DCE	MeOH
CH ₂	NaBH(OAc)₃ HOAc,	HONH ₂ •HCI/NaOMe/
. ✓N	DCE	MeOH
ÇH₂	NaBH(OAc)₃ HOAc,	2 M HONH₂ in MeOH
	DCE	
CH ₂	NaBH₃CN/MeOH/	2 M HONH ₂ in MeOH
N	HOAc	
HN CH2	NaBH(OAc)₃ HOAc,	2 M HONH₂ in MeOH
	DCE	

An alternate method for preparing tertiary amines is by reacting a secondary amine with an alkylating agent in a suitable solvent in the presence of a base. For example, heating a dimethylsulfoxide (DMSO) solution of amine 11 and bromide 40 in the presence of (*i*-Pr)₂NEt yielded tertiary amine 42. Reaction of the tertiary amine 42 with HONH₂•HCl, NaOH in MeOH provides hydroxamate 43. The silyl group can be removed by any method

known to those skilled in the art. For example, the hydroxamate 43 can be treated with an acid, e.g., trifluoroacetic acid, or fluoride to produce hydroxyethyl compound 44.

The hydroxamate compound, or salt thereof, is suitable for preparing pharmaceutical compositions, especially pharmaceutical compositions having deacetylase, especially histone deacetylase, inhibiting properties. Studies with athymic mice demonstrate that the hydroxamate compound causes HDA inhibition and increased histone acetylation *in vivo*, which triggers changes in gene expression that correlate with tumor growth inhibition.

The present invention further includes pharmaceutical compositions comprising a pharmaceutically effective amount of one or more of the above-described compounds as active ingredient. Pharmaceutical compositions according to the invention are suitable for enteral, such as oral or rectal, and parenteral administration to mammals, including man, for the treatment of tumors, alone or in combination with one or more pharmaceutically acceptable carriers.

The hydroxamate compound is useful in the manufacture of pharmaceutical compositions having an effective amount the compound in conjunction or admixture with excipients or carriers suitable for either enteral or parenteral application. Preferred are tablets and gelatin capsules comprising the active ingredient together with (a) diluents; (b) lubricants, (c) binders (tablets); if desired, (d) disintegrants; and/or (e) absorbents, colorants, flavors and sweeteners. Injectable compositions are preferably aqueous isotonic solutions or suspensions, and suppositories are advantageously prepared from fatty emulsions or suspensions. The compositions may be sterilized and/or contain adjuvants,

such as preserving, stabilizing, wetting or emulsifying agents, solution promoters, salts for regulating the osmotic pressure and/or buffers. In addition, the compositions may also contain other therapeutically valuable substances. The compositions are prepared according to conventional mixing, granulating or coating methods, respectively, and contain preferably about 1 to 50% of the active ingredient.

Suitable formulations also include formulations for parenteral administration include aqueous and non-aqueous sterile injection solutions which may contain antioxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the intended recipient; and aqueous and non-aqueous sterile suspensions which may include suspending agents and thickening agents. The formulations may be presented in unit-dose or multi-dose containers, for example, sealed ampules and vials, and may be stored in a freeze-dried (lyophilized) condition requiring only the addition of the sterile liquid carrier, for example, water for injections, immediately prior to use. Extemporaneous injection solutions and suspensions may be prepared from sterile powders, granules and tablets of the kind previously described.

As discussed above, the compounds of the present invention are useful for treating proliferative diseases. A proliferative disease is mainly a tumor disease (or cancer) (and/or any metastases). The inventive compounds are particularly useful for treating a tumor which is a breast cancer, genitourinary cancer, lung cancer, gastrointestinal cancer, epidermoid cancer, melanoma, ovarian cancer, pancreas cancer, neuroblastoma, head and/or neck cancer or bladder cancer, or in a broader sense renal, brain or gastric cancer; in particular (i) a breast tumor; an epidermoid tumor, such as an epidermoid head and/or neck tumor or a mouth tumor; a lung tumor, for example a small cell or non-small cell lung tumor; a gastrointestinal tumor, for example, a colorectal tumor; or a genitourinary tumor, for example, a prostate tumor (especially a hormone-refractory prostate tumor); or (ii) a proliferative disease that is refractory to the treatment with other chemotherapeutics; or (iii) a tumor that is refractory to treatment with other chemotherapeutics due to multidrug resistance.

In a broader sense of the invention, a proliferative disease may furthermore be a hyperproliferative condition such as leukemias, hyperplasias, fibrosis (especially pulmonary, but also other types of fibrosis, such as renal fibrosis), angiogenesis, psoriasis,

atherosclerosis and smooth muscle proliferation in the blood vessels, such as stenosis or restenosis following angioplasty.

Where a tumor, a tumor disease, a carcinoma or a cancer are mentioned, also metastasis in the original organ or tissue and/or in any other location are implied alternatively or in addition, whatever the location of the tumor and/or metastasis.

The compound is selectively toxic or more toxic to rapidly proliferating cells than to normal cells, particularly in human cancer cells, e.g., cancerous tumors, the compound has significant antiproliferative effects and promotes differentiation, e.g., cell cycle arrest and apoptosis. In addition, the hydroxamate compound induces p21, cyclin-CDK interacting protein, which induces either apoptosis or G1 arrest in a variety of cell lines.

The following examples are intended to illustrate the invention and are not to be construed as being limitations thereto.

Example P1

Preparation of *N*-Hydroxy-3-[4-[[[2-(1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide.

4-formylcinnamic acid methylester is produced by adding 4-formylcinnamic acid (25 g, 0.143 mol) in MeOH and HCl (6.7 g, 0.18 mol). The resulting suspension is heated to reflux for 3 hours, cooled and evaporated to dryness. The resulting yellow solid is dissolved in EtOAc, the solution washed with saturated NaHCO₃, dried (MgSO₄) and evaporated to give a pale yellow solid which is used without further purification (25.0 g, 92%). To a solution of tryptamine (16.3 g, 100 mmol) and 4-formylcinnamic acid methylester (19 g, 100 mmol) in dichloroethane, NaBH(OAc)₃ (21 g, 100 mmol) is added. After 4 hours the mixture is diluted with 10% K₂CO₃ solution, the organic phase separated and the aqueous solution extracted with CH₂Cl₂. The combined organic extracts are dried (Na₂SO₄), evaporated and the residue purified by flash chromatography to produce 3-(4-{[2-(1*H*-indol-3-yl)-ethylamino]-methyl}-phenyl)-(2*E*)-2-propenoic acid methyl ester (29 g). A solution of KOH (12.9 g 87%, 0.2 mol) in MeOH (100 mL) is added to a solution of HONH₂•HCl (13.9 g, 0.2 mol) in MeOH (200 mL) and a precipitate results. After 15 minutes the mixture is filtered, the filter cake

washed with MeOH and the filtrate evaporated under vacuum to approximately 75 mL. The mixture is filtered and the volume adjusted to 100 mL with MeOH. The resulting solution 2M HONH₂ is stored under N₂ at -20° C for up to 2 weeks. Then 3-(4-{[2-(1*H*-indol-3-yl)-ethylamino]-methyl}-phenyl)-(2*E*)-2-propenoic acid methyl ester (2.20 g, 6.50 mmol) is added to 2 M HONH₂ in MeOH (30 mL, 60 mmol) followed by a solution of KOH (420 mg, 6.5 mmol) in MeOH (5 mL). After 2 hours dry ice is added to the reaction and the mixture is evaporated to dryness. The residue is dissolved in hot MeOH (20 mL), cooled and stored at -20 °C ovemight. The resulting suspension is filtered, the solids washed with ice cold MeOH and dried under vacuum, producing *N*-Hydroxy-3-[4-[[[2-(1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide (m/z 336 [MH⁺]).

Example P2

Preparation of *N*-Hydroxy-3-[4-[[(2-hydroxyethyl)[2-(1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide

A solution of 3-(4-{[2-(1H-indol-3-yl)-ethylamino]-methyl}-phenyl)-(2E)-2-propenoic acid methyl ester (12.6 g, 37.7 mmol), (2-bromoethoxy)-tert-butyldimethylsilane (12.8 g, 53.6 mmol), (i-Pr)2NEt, (7.42 g, 57.4 mmol) in DMSO (100 mL) is heated to 50° C. After 8 hours the mixture is partitioned with CH₂Cl₂/H₂O. The organic layer is dried (Na₂SO₄) and evaporated. The residue is chromatographed on silica gel to produce 3-[4-([2-(tertbutyldimethylsilanyloxy)-ethyl]-[2-(1H-indol-3-yl)-ethyl]-amino}-methyl)-phenyl]-(2E)-2propenoic acid methyl ester (13.1 g). Following the procedure described for the preparation of the hydroxamate compound in Example P1, 3-[4-({[2-(tert-butyldimethylsilanyloxy)-ethyl]-[2-(1H-indol-3-yl)-ethyl]-amino}-methyl)-phenyl]-(2E)-2-propenoic acid methyl ester (5.4 g, 11 mmol) is converted to N-hydroxy-3-[4-({[2-(tert-butyldimethylsilanyloxy)-ethyl]-[2-(1Hindol-3-yl)-ethyl]-amino}-methyl)-phenyl]-(2E)-2-propenamide (5.1 g,) and used without further purification. The hydroxamic acid (5.0 g, 13.3 mmol) is then dissolved in 95% TFA/H₂O (59 mL) and heated to 40 - 50 °C for 4 hours. The mixture is evaporated and the residue purified by reverse phase HPLC to produce N-Hydroxy-3-[4-[[(2-hydroxyethyl)]2-(1H-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide as the trifluoroacetate salt (m/z 380 [MH+]).

Example P3

Preparation of N-hydroxy-3-[4-[[[2-(2-methyl-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide.

A suspension of LiAlH₄ (17 g, 445 mmol) in dry THF (1000 mL) is cooled to 0 °C and 2methylindole-3-glyoxylamide (30 g, 148 mmol) is added in portions over 30 min. The mixture is stirred at room temperature for 30 min. and then maintained at reflux for 3 h. The reaction is cooled to 0 °C and treated with H₂O (17ml), 15% NaOH (aq., 17ml) and H₂O (51ml). The mixture is treated with MgSO₄, filtered and the filtrate evaporated to give 2-methyltryptamine which is dissolved in MeOH. Methyl 4-formylcinnamate (16.9 g, 88.8 mmol) is added to the solution, followed by NaBH₃CN (8.4 g) and AcOH (1 equiv.). After 1h the reaction is diluted with NaHCO₃ (aq.) and extracted with EtOAc. The organic extracts are dried (MgSO₄), filtered and evaporated. The residue is purified by chromatography to give 3-(4-{[2-(2methyl-1H-indol-3-yl)-ethylamino]-methyl}-phenyl)-(2E)-2-propenoic acid methyl ester. The ester is dissolved in MeOH, 1.0 M HCl/dioxane (1 - 1.5 equiv.) is added followed by Et₂O. The resulting precipitate is filtered and the solid washed with Et₂O and dried thoroughly to give 3-(4-{[2-(2-methyl-1*H*-indol-3-yl)-ethylamino]-methyl}-phenyl)-(2*E*)-2-propenoic acid methyl ester hydrochloride. 1.0 M NaOH (aq., 85 mL) is added to an ice cold solution of the methyl ester hydrochloride (14.9 g, 38.6 mmol) and HONH₂ (50% aq. solution, 24.0 mL, ca. 391.2 mmol). After 6 h, the ice cold solution is diluted with H₂O and NH₄Cl (aq., 0.86 M, 100 mL). The resulting precipitate is filtered, washed with H₂O and dried to afford N-hydroxy-3-[4-[[[2-(2-methyl-1 H-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide (m/z 350 $[MH^{\dagger}]$).

Examples 1-265

The following compounds are prepared by methods analogous to those disclosed in Examples P1, P2 and P3:

Example	STRUCTURE	m/z (MH*)
1		426

Example	STRUCTURE	m/z (MH ⁺)
2	Jan Ott	
3	Thomas and the second s	·
4	N-OH	325
5	Ů p oH	
6	· CHOH	
7		·
8 .	HAN OH	465

Example	STRUCTURE	m/z (MH†)
9	N N N N N N N N N N N N N N N N N N N	·
10	ни	
11	CH CH	
12	ino Hyo	420
13	HN	420

Example	STRUCTURE	m/z (MH ⁺)
14	J J J J J J J J J J J J J J J J J J J	
15	OH OH	465
16	NOH OH	385
17	HO H	550
18	Non Non	432
19		366

Example	STRUCTURE	m/z (MH ⁺)
20	HOH NAME OF THE OWN	350
21	но он мага и он	
22	D HO HO	442
23		338
24	S NH	464
25	рон Пон	541

Example	STRUCTURE	m/z (MH†)
26	OH NO DH	
27	The state of the s	
28	The state of the s	417
29	L L L L L L L L L L L L L L L L L L L	
30	H J H OH	
31	D OH	380
32	J. OH	436
33	The state of the s	

Example	STRUCTURE	m/z (MH ⁺)
34		493
35	N N N N N N N N N N N N N N N N N N N	477
36		586
37		513
38		378
39		408

Example	STRUCTURE	m/z (MH ⁺)
40	H H H	449
41	NH PH	438
42		452
43		507
44		565

Example	STRUCTURE	m/z (MH ⁺)
45		
46	J. OH J. OH	
47	Д ОН Д ОН	
48	CH CH	
49	The state of the s	
50	The state of the s	·

Example	STRUCTURE	m/z (MH⁺)
51		470
52	OH OH	
53	OH OH	548
54	NH OH	623
55		456
56	NH OH	478

Example	STRUCTURE	m/z (MH⁺)
57	OH JUNE	394
58		422
59	OH OH	479
60	H-OH	603
61		477

Example	STRUCTURE	m/z (MH ⁺)
62	NOH OH	539
63	HIN OH	523
64	H C C C C C C C C C C C C C C C C C C C	·
65	он — — — — — — — — — — — — — — — — — — —	
66	The state of the s	
67	The state of the s	

Example	STRUCTURE	m/z (MH⁺)
68	I NOH	539
69		495
70	De la companya de la	
71	H ₂ N OH	379
72		478

Example	STRUCTURE	m/z (MH ⁺)
73		462
74		378
75	С В С В С В С В С В С В С В С В С В С В	
76		493
77		503
78	L COH	350

Example	STRUCTURE	m/z (MH†)
79		549
80	D D D D D D D D D D D D D D D D D D D	471
81	O No	350
82		418
83	OH OH	486

Example	STRUCTURE	m/z (MH ⁺)
84	F F OH OH	524
85	OH OH	424
86		364
87		440
88		420

Example	STRUCTURE	m/z (MH†)
89		390
90		
91	H OH	
92	H OH	484
93	THE STATE OF THE S	498
94	J J J J J J J J J J J J J J J J J J J	490

Example	STRUCTURE	m/z (MH ⁺)
95	NON DON DON DON DON DON DON DON DON DON	
96	HN OH	475
97	HN N OH	525
98	H OH	422
99	TH OH	528

Example	STRUCTURE	m/z (MH⁺)
100	P P P P P P P P P P P P P P P P P P P	448
101	HA OH	437
102		451
103	H OH	505

Example	STRUCTURE	m/z (MH⁺)
104	HN OH	519
105	HO SHOW	514
106	HNOH	507
107	OH OH	626

Example	STRUCTURE	m/z (MH ⁺)
108	HBN OH	499
109	HN OH	
110	ны по	
111	O C C C C C C C C C C C C C C C C C C C	429
112		464
113		432

Example	STRUCTURE	m/z (MH ⁺)
114	OH OH	422
115		390
116	OH OH	501
117	HIN OH	484
118	NH HO	·

Example	STRUCTURE	m/z (MH ⁺)
119	HN OH	587
120	HN OH	602
121		539
122	HIN CH	
123	The state of the s	528

Example	STRUCTURE	m/z (MH⁺)
124	THO OH	487
125	O=S=O N OH	
126	AND	556
127	ONE O.	
128	HN OH	
129	To the Ho	552

Example	STRUCTURE	m/z (MH⁺)
130	OH OH SEEO	519
131	The state of the s	450
132	H OH	464
133	OH OH	558
134	NOH OH	533

Example	STRUCTURE	m/z (MH ⁺)
135	O O O O O O O O O O O O O O O O O O O	
136		527
137	N OH	381
138		364
139	HDZ NOH	
140	DE LA COMPANSION DE LA	448

Example	STRUCTURE	m/z (MH⁺)
141	OH OH	558
142	HO H	
143	DH HZ DH	427
144		,
145	O J N O OH	432
146	HO-H	384
147		354

Example	STRUCTURE	m/z (MH⁺)
148	O S O O N O O O O O O O O O O O O O O O	
149		
150	O=S=O NOH	
151	O O O O O O O O O O O O O O O O O O O	
152	HN NOH	
153	N OH	

Example	STRUCTURE	m/z (MH⁺)
154		350
155	HO HO HO	366
156		408
157	NH OH	322
158	HINDHOH	364
159		364
160	The second secon	378

Example	STRUCTURE	m/z (MH⁺)
161	J. J. J. P. OH	350
162		463
163	O D D D D D D D D D D D D D D D D D D D	
164	The state of the s	381
165	от распорация и по	463
166		476
167	HALL OH	

Example	STRUCTURE	m/z (MH ⁺)
168	OH OH	
169	NH OH	
170	J N N N OH	368
171	OH OH	493
172	HIN OH	527
173	HD N OH	515

Example	STRUCTURE	m/z (MH ⁺)
174	N N N N N N N N N N N N N N N N N N N	323
175	HN ON OH	540
176	у проставля про	441
177	л О П он	276
178	I OH	
179	HOH	455
180	PIN OH	

Example	STRUCTURE	m/z (MH⁺)
181	De la contraction de la contra	336
182		347
183	2 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 - 3 -	447
184	N N N OH	
185	C C C C C C C C C C C C C C C C C C C	420
186		424

Example	STRUCTURE	m/z (MH⁺)
187	THE OH	422
188	The state of the s	
189	F OH OH	398
190	HN OH	418
191		350
192	HO OH	
193		352

Example	STRUCTURE	m/z (MH ⁺)
194	OH OH	499
195	он он	408
196	HO J H OH	394
197	J. J. OH	499
198		

Example	STRUCTURE	m/z (MH ⁺)
199	O O O O O O O O O O O O O O O O O O O	
200	H H H OH	350
201	он он он	
202	он Он	
203	H ₂ M NH HO	
204	H ₂ N NH OH	365

Example	STRUCTURE	m/z (MH ⁺)
205		465
206	NH ₂ OH	
207	OH OH OH	410
208	HO HO OH	410
209	р н	·
210	он он	366

Example	STRUCTURE	m/z (MH⁺)
211		352
212	HO I NO OH	
213	р — — — — — — — — — — — — — — — — — — —	368
214		338
215	A NOW NOW	356
216	H OH	408
217	N OH	368

Example	STRUCTURE	m/z (MH ⁺)
218	NH HO	396
219	THE STATE OF THE S	
220		342
221	S H OH	392
222	L L OH	412
223	Charles Charle	337
224	HIN THOM	337

Example	STRUCTURE	m/z (MH ⁺)
225	HO OH	456
226	HOH OH	364
227	HO ZH	481
228	NH2 OH	355
229	C NH H OH	312
230	HO NH HO	424

Example	STRUCTURE	m/z (MH ⁺)
231		
232	OH NOH	351
233		392
234	P P P	
235	OH OH	
236	NH OH	322
237	I I I OH	

Example	STRUCTURE	m/z (MH ⁺)
238	OH OH OH	366
239	D P P P	
240	THE STATE OF THE S	368
241	P P P P P P P P P P P P P P P P P P P	
242	D OH	. 406
243	NH OH	398
244	NE N	442

Example	STRUCTURE	m/z (MH⁺)
245	F = 2	350
246	NH OH	364
247	NH OH	402
248	1	418
249	OH OH	. 364
250	P P P P P P P P P P P P P P P P P P P	

Example	STRUCTURE	m/z (MH ⁺)
251	OH OH	408
252	H OH	
253	THO OH	
254	HAN THE THE PART OF THE PART O	413
255	OH OH	405
256		

Example	STRUCTURE	m/z (MH⁺)
257	DE CONTROLL OF THE CONTROL OF THE CO	394
258	Charles on the one	390
259	DH OH	434
260		386
261	HO J H OH	368
262	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	412

Example	STRUCTURE	m/z (MH ⁺)
263	J H OH	406
264	C C C C C C C C C C C C C C C C C C C	
265	I OH	378

The compounds of Examples 1-265 show an HDA enzyme IC $_{50}$ in the range from about 0.005 to about 0.5 μ M.

Example B1

Cell lines H1299 (human lung carcinoma cell) and HCT116 (colon tumor cell) are obtained from the American Type Culture Collection, Rockville, MD. The cell lines are free of *Mycoplasma* contamination (Rapid Detection System by Gen-Probe, Inc., San Diego, CA) and viral contamination (MAP testing by MA BioServices, Inc., Rockville, MD). The cell lines are propagated and expanded in RPMI 1640 medium containing 10% heat-inactivated FBS (Life Technologies, Grand Island, NY). Cell expansions for implantation are performed in cell factories (NUNC, purchased from Fisher Scientific, Springfield, NJ). Cells are harvested at 50-90% confluency, washed once with HBSS (Hank's Balanced Salt Solution) containing 10% FBS, and suspended in 100% HBSS.

Cell proliferation is measured with a commercial MTS kit (Promega, Madision, Wis.) assay using an adaptation of published procedures, for example, that disclosed in Feasibility of drug screening with panels of human tumor cell lines using a microculture tetrazolium assay, Alley MC, et al., Cancer Res. 1988; 48:589-601. Cells are plated in 96-well tissue culture dishes, with top and bottom rows left empty. H1299 and HCT116 cells

are suspended in complete media at a density of 5.3×10^3 and 3.6×10^3 cell/mL, respectively, and 190 μ l are added per well. Each cell line is added to one half of the plate. Complete medium (200 μ L) is added to the top and bottom rows. Twenty-four hours later, 10 μ l of MTS solution is added to one of the plates to determine the activity at the time of compound addition (T_0). The plate is incubated at 37 °C for 4 hours and the OD₄₉₀ is measured on a Molecular Devices Thermomax at 490 nm using the Softmax program. The T_0 plate serves as a reference for initial activity at the beginning of the experiment.

Five serial dilutions (1:4) of each compound are made in a 96-deep well plate with the highest concentrations on the edge of plate. Two cell lines are tested with two compounds per plate. Ten microliters of each of the five dilutions are added in triplicate and complete medium alone is added to columns six and seven. The plates are incubated at 37 °C for 72 hours. The MTS solution is added (as for the T₀ plate) and read four hours later.

In order to analyze the data, the average background value (media alone) is subtracted from each experimental well; the triplicate values are averaged for each compound dilution. The following formulas are used to calculate percent growth.

If $X > T_0$, % Growth = $((X-T_0)/(GC - T_0)) \times 100$

If $X < T_0$, % Growth = $(X-T_0)/T_0$) x 100

in which To = (average value of cell viability at time 0) - background

GC = average value of untreated cells (in triplicate) - background

X = average value of compound treated cells (in triplicate) - background

The "% Growth" is plotted against compound concentration and used to calculate IC₅₀s employing the linear regression techniques between data points to predict the concentration of compounds at 50% inhibition.

Lactate salts of N-hydroxy-3-[4-[[[2-(1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide (CMD1), N-hydroxy-3-[4-[[(2-hydroxyethyl)[2-(1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide (CMD2), N-hydroxy-3-[4-[[[2-(5-methoxy-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide (CMD3), N-hydroxy-3-[4-[[[2-(5-fluoro-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide (CMD4), N-hydroxy-3-[4-[[[2-(benzofur-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide (CMD5) having a purity of higher than 95% are dissolved in pure dimethylsulfoxide (DMSO) to create a stock solution. The stock solution is diluted with 5% dextrose injection, USP, just prior to dosing. In addition, N-(2-aminophenyl)-4-[N-pyridin-3-yl)methoxycarbonylaminomethyl]benzamide is synthesized in accordance with Example 48 of EP 0 847 992 and used as a control compound (CMDC). Inhibition of cell growth in monolayer for 72 hours of compound treatment is measured in

triplicate experiments and used to derive the IC_{50} by MTS assay. The results are shown in Table B1.

Table B1

Monolayer Growth IC50 (µM)

Compound	<u>H1299</u>	HCT116
CMD1	0.40	0.03
CMD2	0.15	0.01
CMD3	0.58	0.03
CMD4	0.28	0.03
CMD5	0.18	0.03
CMDC	6.8	0.67

The results show that the hydroxamate compounds of the present invention are highly active in inhibition of tumor cell growth. In addition to the above results, it has been observed that the compounds selectively inhibited tumor cells while showing minimal inhibition activities in non-tumorous cells.

The cells treated with the hydroxamate compounds are also tested for the induction of p21 promoter, which is a key mediator of G1 arrest and differentiation. The hydroxamate compounds activate the p21 promoter to a readily detectable level at a concentration within two-fold of their respective IC₅₀ for monolayer cell growth inhibition in H1299. Without being bound by any particular theory, the correlation appears to demonstrate that HDA inhibition leads to transcriptional activation of genes that inhibit tumor cell proliferation.

Example B2

HDA is partially purified from H1299, human non-small cell lung carcinoma cells (obtained from American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD 20852, USA). Cells are grown to 70-80% confluence in RPMI media in the presence of 10% FCS, harvested and lysed by sonication. The lysate is centrifuged at 23, 420g for 10-15 min, the supernatant is applied to a Hiload 26/10 High performance Q-sepharose column (Amersham Pharmacia Biotech), and equilibrated with a buffer containing 20 mM Tris

pH8.0, 1 mM EDTA, 10 mM NH₄Cl₂, 1 mM β-Mercaptoethanol, 5% glycerol, 2 µg/mL aprotinin, 1 µg/mL leupeptin, and 400 mM PMSF. Proteins are eluted in 4mL aliquotes with a linear gradient from 0-500 mM NaCl in the above buffer at a flow rate of 2.5 mL/min. Each preparation of partially purified HDA enzyme is titrated to determine the optimal amount needed to obtain a signal to noise ratio of at least 5 to 1. Generally, 20-30 µl of partially purified HDA (5-10 mg protein/mL) is mixed with 2 μ L of compound solution in DMSO in a deep well titer plate (Beckman). The compounds are serially diluted in DMSO to generate stocks at 20-fold of the assay concentrations. Final concentrations of compounds in the assay are 10 μM, 2 μM, 400 nM, 80 nM, and 16 nM with the final percentage of DMSO in each enzyme reaction equaling 0.1%. Each concentration of compound is assayed in duplicate. The substrate used in the reaction is a peptide of amino acid sequence, SGRGKGGKGLGKGGAKRHRKVLRD, corresponding to the twenty-four N-terminal amino acids of human histone H4, biotinylated at the N-terminus and penta-acetylated, at each lysine residue with $^3\text{H-acetate}$. To initiate the reaction, the substrate is diluted in 10 μL of Buffer A (100 mM Tris pH 8.0, 2 mM EDTA), added to the enzyme mixture and collected at the bottom of the deep well plate by centrifugation for 5 minutes at 1500 rpm. Following centrifugation, the mixture is incubated at 37 °C for 1.5 hr. The reaction is stopped by the addition of 20 µL of the Stop Buffer (0.5N HCl, 0.08M Acetic Acid). At this point, the assay proceeds to the robotic extraction phase or is frozen for several days at -80 °C.

The extraction of enzymatically cleaved ³H-acetate groups from the reaction mixture is achieved with the solvent TBME (t-butyl methyl ether) using the Tomtec Quadra 96 workstation. A program is written to add 200 µL of TBME to a 96 "deep well" plate. The workstation is programmed to aspirate 50 µL of air followed by 200 µL of TBME and finally another 25 µL of air, which is dispensed into the each well of the plate. The contents of the deep well were mixed thoroughly by pipetting 160 µL up and down 10 times. Before addition of TBME to the reaction mixture, it is necessary to "pre-wet" the pipette tips with TBME to prevent the solvent from dripping during the transfer to the deep well plate. The organic and aqueous phases in the deep well are separated by centrifugation at 1500 rpm for 5 min. Opti-Phase Supermix liquid scintillation cocktail (200 µL) (Wallac) is added to each well of the 96-well Trilux plate (Wallac). The deep well and Trilux plates are placed back on the workstation programmed to aspirate 25 µL of air into the pipette tips followed by 100 µL of the upper TBME phase and transfer it into the Trilux plate. The solutions are mixed by pipetting and expelling 50 µL, five times, within the same well. The Trilux plate is

covered with clear film and read on a 1450 MicroBeta Trilux liquid scintillation and luminescence counter (Wallac) with a color/chemical quench and dpm correction.

In order to determine the IC₅₀ values, the data are analyzed on a spreadsheet. The analysis requires a correction for the background luminescence that is accomplished by subtracting the dpm values of wells without ³H substrate from the experimental wells. The corrected dpm values along with the concentrations of the compounds are used to calculate IC₅₀ using the user-defined spline function. This function utilizes linear regression techniques between data points to calculate the concentration of compounds that produced 50% inhibition. The results are shown in Table B2.

Table B2

Compound	HDA Enzyme Activity IC ₅₀ (μM)
CMD1	0.032
CMD2	0.063
CMD3	0.014
CMD4	0.014
CMD5	0.016
CMDC	> 10

Example B3

The A549 non-small cell lung human tumor cell line is purchased from the American Type Culture Collection, Rockville, MD. The cell line is free of *Mycoplasma* contamination (Rapid Detection System by Gen-Probe, Inc., San Diego, CA) and viral contamination (MAP testing by MA BioServices, Inc., Rockville, MD). The cell line is propagated and expanded in RPMI 1640 medium containing 10% heat-inactivated FBS (Life Technologies, Grand Island, NY). Cell expansions for implantation are performed in cell factories (NUNC, purchased from Fisher Scientific, Springfield, NJ). Cells are harvested at 50-90% confluency, washed once with HBSS containing 10% FBS, and suspended in 100% HBSS.

Outbred athymic (nu/nu) female mice ("Hsd:Athymic Nude-nu" from Harlan Sprague Dawley, Indianapolis, IN) are anesthetized with Metofane (Mallinckrodt Veterinary, Inc.,

Mundelein, IL), and 100 μ L of the cell suspension containing $1x10^7$ cells is injected subcutaneously into the right axillary (lateral) region of each animal. Tumors are allowed to grow for about 20 days until a volume of ~100 mm³ is achieved. At this point, mice bearing tumors with acceptable morphology and size are sorted into groups of eight for the study. The sorting process produces groups balanced with respect to mean and range of tumor size. Antitumor activity is expressed as % T/C, comparing differences in tumor volumes for treatment group (T) to vehicle control group (C). Regressions are calculated using the formula: $(1-T/T_0) \times 100\%$, where T is the tumor volume for the treatment group at the end of the experiment, and T_0 is the tumor volume at the beginning of the experiment.

CMD1 is administered intravenously, once daily 5x/week for three weeks, at doses of 10, 25, 50, or 100 mg/kg. The final DMSO concentration is 10%. Each test group has eight mice. Tumors are measured, and individual animal body weights recorded. Table B3 shows the results on the 41st day.

Table B3

		Δ MEAN		Δ%
	DOSE	TUMOR VOLUME ¹¹		BODY WEIGHT ²
COMPOUND	<u>(mg/kg)</u>	(mm³ ± SEM⁴³)	% T/C	(% ± SEM*3)
10% DMSO/D5W ^{*4}	-	376 ± 55	-	+11.9 ± 0.2
CMD1	10	121 ± 27	32	+ 1.3 ± 0.3
CMD1	25	77 ± 32	20	- 0.9 ± 0.3
CMD1	50	57 ± 10	15	- 0.4 ± 0.3
CMD1	100	28 ± 25	7	+ 0.4 ± 0.3

Note: *1. Difference in mean tumor volume for a group of animals at the end of the experiment minus mean tumor volume at the beginning.

- *2. Difference in body weight for a group of animals at the end of the experiment minus mean tumor volume at the beginning.
 - *3. Standard error of the mean.
 - *4.5% dextrose injection, USP.

Example B4

Example B3 repeated except CMD2 is used. Table B4 shows the results.

Table B4

		Δ ΜΕΑΝ		Δ%
	DOSE	TUMOR VOLUME		BODY WEIGHT
COMPOUND	(mg/kg)	(mm³ ± SEM)	% T/C	(% ± SEM)
10% DMSO/D5W	-	135 ± 43	-	+ 6.7 ± 1.1
CMD2	25	37 ± 16	27	- 4.2 ± 2.5
CMD2	50	29 ± 15	21	- 2.9 ± 1.5

Example B5

Example B3 is repeated except the HCT116 colon tumor cell line is used in place of the A549 cell line. The HCT116 cell line is also obtained from American Type Culture Collection, Rockville, MD, and the cell line is free of *Mycoplasma* contamination and viral contamination. The results are recorded on the 34th day and are shown in Table B5.

Table B5

		Δ MEAN		Δ%
	DOSE	TUMOR VOLUME	-	BODY WEIGHT
COMPOUND	(mg/kg)	(mm³ ± SEM)	.% T/C	(% ± SEM)
10% DMSO/D5W	-	759 ± 108	-	- 0.4 ± 0.4
CMD1	50 ^{*10}	186 ± 40	25	- 7.4 ± 0.8
CMD1	100	140 ± 38	18	- 3.2 ± 0.4
Note: *10. Seven mid	ce are tested	d in this aroun		

Example B6

Example B4 is repeated except the HCT116 colon tumor cell line is used in place of the A549 cell line. The HCT116 is also obtained from American Type Culture Collection, Rockville, MD, and the cell line is free of *Mycoplasma* contamination and viral contamination. The results are recorded on the 34th day and are shown in Table B6.

Table B6

		Δ MEAN		Δ%
	DOSE	TUMOR VOLUME		BODY WEIGHT
COMPOUND	<u>(mg/kg)</u>	(mm³ ± SEM)	% T/C	<u>(% ± SEM)</u>
10% DMSO/D5W	-	759 ± 108	-	- 0.4 ± 0.4
CMD2	10	422 ± 75	56	- 10.2 ± 0.5
CMD2	25	305 ± 47	40	- 7.0 ± 0.2
CMD2	50	97 ± 30	13	- 7.3 ± 0.3
CMD2	100	132 ± 30	17	- 9.4 ± 0.4

Example B7

Annexin V binding was used as a marker for the early stages of apoptosis. A549, HCT116 and Normal Dermal Human Fibroblasts (NDHF) cells are treated separately with four compounds (CMD1, CMD2, CMD3 and CMD4) for 24 or 48 hours, stained with annexin V and compared to cells treated similarly with vehicle (DMSO). Cells are examined by fluorescence microscopy. Those undergoing apoptosis exhibit green fluorescent membrane staining. Viability is assessed by the counterstain, propidium iodide. Cells detected by red fluorescence are not viable. A small percentage of A549 and the majority of HCT116 cells exhibit cell surface staining with annexin V after 24 hour exposure to each of the four compounds. After 48 hour treatment, the majority of the A549 and HCT116 stain with annexin V and/or propidium iodide indicating that the compounds induce apoptotic cell death. In contrast, NDHF cells do not show noticeable annexin V staining after 24 hour exposure and limited annexin V staining with CMD3 after 48 hour. These data show that

- 81 -

NDHF cells predominantly underwent non-lethal growth arrest upon compound treatment, consistent with the cell cycle profile.

The staining results demonstrate that the hydroxamate compounds of the present invention cause tumor cells to die by apoptosis, while causing normal fibroblast to predominantly undergo cell cycle arrest, clearly demonstrating the selective efficacy of the present compounds.

What is claimed is:

1. A compound of the formula I

HO
$$R_1$$
 R_2 R_3 R_4 R_5 R_5 R_5

wherein

R₁ is H, halo, or a straight chain C₁-C₆ alkyl;

R₂ is selected from H, C₁-C₁₀ alkyl, C₄ – C₉ cycloalkyl, C₄ – C₉ heterocycloalkyl, C₄ – C₉ heterocycloalkylalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, -(CH₂)_nOC(O)R₆, amino acyl, HON-C(O)-CH=C(R₁)-aryl-alkyl- and -(CH₂)_nR₇;

R₃ and R₄ are the same or different and independently H, C₁-C₆ alkyl, acyl or acylamino, or R₃ and R₄ together with the carbon to which they are bound represent C=O, C=S, or C=NR₈, or R₂ together with the nitrogen to which it is bound and R₃ together with the carbon to which it is bound can form a C₄ - C₉ heterocycloalkyl, a heteroaryl, a polyheteroaryl, a non-aromatic polyheterocycle, or a mixed aryl and non-aryl polyheterocycle ring;

R₅ is selected from H, C₁-C₆ alkyl, C₄ – C₉ cycloalkyl, C₄ – C₉ heterocycloalkyl, acyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, aromatic polycycle, non-aromatic polycycle, mixed aryl and non-aryl polycycle, polyheteroaryl, non-aromatic polyheterocycle, and mixed aryl and non-aryl polyheterocycle;

 n_1 , n_2 and n_3 are the same or different and independently selected from 0 – 6, when n_1 is 1-6, each carbon atom can be optionally and independently substituted with R_3 and/or R_4 ;

X and Y are the same or different and independently selected from H, halo, C₁-C₄ alkyl, NO₂, C(O)R₁, OR₉, SR₉, CN, and NR₁₀R₁₁;

- R₆ is selected from H, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, OR₁₂, and NR₁₃R₁₄;
- R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{16}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_6$;
- R_8 is selected from H, OR_{15} , $NR_{13}R_{14}$, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_9 is selected from $C_1 C_4$ alkyl and C(O)-alkyl;
- R₁₀ and R₁₁ are the same or different and independently selected from H, C₁-C₄ alkyl, and -C(O)-alkyl;
- R_{12} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, C_4 C_9 heterocycloalkylalkyl, aryl, mixed aryl and non-aryl polycycle, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, amino acyl, or R_{13} and R_{14} together with the nitrogen to which they are bound are C_4 C_9 heterocycloalkyl, heteroaryl, polyheteroaryl, non-aromatic polyheterocycle or mixed aryl and non-aryl polyheterocycle;
- R_{15} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{16} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, polyheteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{17} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, aromatic polycycle, heteroaryl, arylalkyl, heteroarylalkyl, polyheteroaryl and $NR_{13}R_{14}$; m is an integer selected from 0 to 6; and
- Z is selected from O, NR₁₃, S and S(O);
- or a pharmaceutically acceptable salt thereof.
- 2. A compound of claim 1 wherein each of R_1 , X, Y, R_3 , and R_4 is H.
- 3. A compound of claim 2 wherein one of n_2 and n_3 is zero and the other is 1.
- 4. A compound of claim 3 wherein R₂ is H or -CH₂-CH₂-OH.
- 5. A compound of claim 1 of the formula la

wherein

n₄ is 0-3,

 R_2 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH_2)_nC(O) R_6 , amino acyl and -(CH_2)_n R_7 ;

R₅' is heteroaryl, heteroarylalkyl, an aromatic polycycle, a non-aromatic polycycle, a mixed aryl and non-aryl polycycle, polyheteroaryl, or a mixed aryl and non-aryl polyheterocycle,

or a pharmaceutically acceptable salt thereof.

6. A compound of claim 1 of the formula la

HO
$$R_{5}$$
 (Ia)

wherein

n₄ is 0-3,

 R_2 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, amino acyl and -(CH₂)_nR₇;

R₅' is aryl, arylalkyl, an aromatic polycycle, a non-aromatic polycycle or a mixed aryl and non-aryl polycycle,

or a pharmaceutically acceptable salt thereof.

7. A compound of claim 6 wherein R_5 is anyl or anylalkyl.

- 8. A compound of claim 7 wherein R_5 ' is p-fluorophenyl, p-chlorophenyl, p-O-C₁-C₄-alkylphenyl, p-C₁-C₄-alkylphenyl, benzyl, ortho, meta or para-fluorobenzyl, or ortho, meta or para-chlorobenzyl, or ortho, meta or para mono, di or tri-O-C₁-C₄-alkylbenzyl.
- 9. A compound of claim 1 of the formula lb

wherein

 R_2 ' is selected from H, C_1 - C_6 alkyl, C_4 - C_6 cycloalkyl, cycloalkylalkyl, -(CH_2)₂₋₄ OR_{21} where R_{21} is H, methyl, ethyl, propyl, or isopropyl, and

 R_5 " is unsubstituted or substituted 1*H*-indol-3-yl, benzofuran-3-yl or quinolin-3-yl, or a pharmaceutically acceptable salt thereof.

- 10. A compound of claim 9 wherein R_5 " is substituted 1*H*-indol-3-yl or substituted benzofuran-3-yl.
- 11. A compound of claim 1 of the formula lc

HO N
$$R_1$$
 R_{18} R_{18} R_{19} R_{19}

wherein

the ring containing Z_1 is aromatic or non-aromatic which non-aromatic rings are saturated or unsaturated,

Z₁ is O, S or N-R₂₀;

 R_{18} is H, halo, C_1 - C_6 alkyl, C_3 - C_7 cycloalkyl, aryl, or heteroaryl;

 R_{20} is H, C_1 - C_6 alkyl, C_1 - C_6 alkyl- C_3 - C_9 cycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, acyl or sulfonyl;

A₁ is 1, 2 or 3 substituents which are independently H, C₁-C-₆alkyl, -OR₁₉, halo, alkylamino, aminoalkyl, halo, or heteroarylalkyl;

 R_2 is selected from H, C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, amino acyl and -(CH₂)_nR₇;

R₁₉ is selected from H, C₁-C₆alkyl, C₄-C₉cycloalkyl, C₄-C₉heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;

v is 0, 1 or 2,

p is 0-3, and

q is 1-5 and r is 0 or

q is 0 and r is 1-5,

or a pharmaceutically acceptable salt thereof.

- 12. A compound of claim 11 wherein Z₁ is N-R₂₀.
- 13. A compound of claim 11 wherein R₂ is H or -CH₂-CH₂-OH and the sum of q and r is 1.
- 14. A compound of claim 1 of the formula ld

HO
$$R_1$$
 R_{18} R_{18} R_{19} R

wherein

Z₁ is O, S or N-R₂₀,

R₁₈ is H, halo, C₁-C₆alkyl, C₃-C₇cycloalkyl, unsubstituted phenyl, substituted phenyl, or heteroaryl,

 R_{20} is H, C_1 - C_6 alkyl, C_1 - C_6 alkyl- C_3 - C_9 cycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, acyl or sulfonyl;

A₁ is 1, 2 or 3 substituents which are independently H, C_1 - C_{-6} alkyl, - OR_{19} , or halo, R_{19} is selected from H, C_1 - C_6 alkyl, C_4 - C_9 cycloalkyl, C_4 - C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and - $(CH_2CH=CH(CH_3)(CH_2))_{1-3}H$;

p is 0-3, and

q is 1-5 and r is 0 or

q is 0 and r is 1-5,

or a pharmaceutically acceptable salt thereof.

- 15. A compound of claim 14 wherein R₂ is H or –CH₂-CH₂-OH and the sum of q and r is 1.
- 16. A compound of claim 11 of the formula le

HO N
$$R_1$$
 R_2 R_3 R_4 R_{18} $N-R_{20}$ (Ie)

or a pharmaceutically acceptable salt thereof.

- 17. A compound of claim 16 wherein R_{18} is H, fluoro, chloro, bromo, C_1 - C_4 alkyl, C_3 - C_7 cycloalkyl, phenyl or heteroaryl.
- 18. A compound of claim 16 wherein R_2 is H, or -(CH₂)_pCH₂OH and wherein p is 1-3.
- 19. A compound of claim 18 wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3.

- 20. A compound of claim 16 wherein R₁₈ is H, methyl, ethyl, t-butyl, trifluoromethyl, cyclohexyl, phenyl, 4-methoxyphenyl, 4-trifluoromethylphenyl, 2-furanyl, 2-thiophenyl, or 2-, 3- or 4-pyridyl.
- 21. A compound of claim 20 wherein R₂ is H, or -(CH₂)_pCH₂OH.
- 22. A compound of claim 21 wherein p is 1-3.
- 23. A compound of claim 22 wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3.
- 24. A compound of claim 23 wherein R₂ is H or -CH₂-CH₂-OH and the sum of q and r is 1.
- 25. A compound of claim 16 wherein R₂₀ is H or C₁-C₆alkyl.
- 26. A compound of claim 16 selected from the group consisting of N-hydroxy-3-[4-[[(2-hydroxyethyl)]2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, N-hydroxy-3-[4-[[[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide and N-hydroxy-3-[4-[[[2-(2-methyl-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 27. A compound of claim 26 which is N-hydroxy-3-[4-[[(2-hydroxyethyl)[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 28. A compound of claim 1 of the formula If

HO N
$$R_{18}$$
 (If)

or a pharmaceutically acceptable salt thereof.

- 29. A compound of claim 28 wherein R₂ is H or -(CH₂)_pCH₂OH and p is 1-3.
- 30. A compound of claim 29 wherein R_1 is H and X and Y are each H, and wherein q is 1-3 and r is 0 or wherein q is 0 and r is 1-3.
- 31. A compound of claim 30 wherein R₂ is H or –CH₂-CH₂-OH and the sum of q and r is 1.
- 32. A compound of claim 28 which is N-hydroxy-3-[4-[[[2-(benzofur-3-yl)-ethyl]-amino]methyl]phenyl]-2E-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 33. A pharmaceutical composition comprising a pharmaceutically effective amount of a compound of formula I

wherein

R₁ is H, halo, or a straight chain C₁-C₈ alkyl;

- R_2 is selected from H, C_1 - C_{10} alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkylalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₆, -(CH₂)_nOC(O)R₈, amino acyl, HON-C(O)-CH=C(R₁)-aryl-alkyl- and -(CH₂)_nR₇;
- R₃ and R₄ are the same or different and independently H, C₁-C₆ alkyl, acyl or acylamino, or R₃ and R₄ together with the carbon to which they are bound represent C=O, C=S, or C=NR₈, or R₂ together with the nitrogen to which it is bound and R₃ together with the carbon to which it is bound can form a C₄ C₉ heterocycloalkyl, a heteroaryl, a polyheteroaryl, a non-aromatic polyheterocycle, or a mixed aryl and non-aryl polyheterocycle ring;

- R₅ is selected from H, C₁-C₆ alkyl, C₄ − C₉ cycloalkyl, C₄ − C₉ heterocycloalkyl, acyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, aromatic polycycle, non-aromatic polycycle, mixed aryl and non-aryl polycycle, polyheteroaryl, non-aromatic polyheterocycle, and mixed aryl and non-aryl polyheterocycle;
- n_1 , n_2 and n_3 are the same or different and independently selected from 0 6, when n_1 is 1-6, each carbon atom can be optionally and independently substituted with R_3 and/or R_4 ;
- X and Y are the same or different and independently selected from H, halo, C₁-C₄ alkyl, NO₂, C(O)R₁, OR₉, SR₉, CN, and NR₁₀R₁₁;
- R₆ is selected from H, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, OR₁₂, and NR₁₃R₁₄;
- R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{16}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_6$;
- R₈ is selected from H, OR₁₅, NR₁₃R₁₄, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_9 is selected from $C_1 C_4$ alkyl and C(O)-alkyl;
- R₁₀ and R₁₁ are the same or different and independently selected from H, C₁-C₄ alkyl, and -C(O)-alkyl;
- R_{12} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, C_4 C_9 heterocycloalkylalkyl, aryl, mixed aryl and non-aryl polycycle, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, amino acyl, or R_{13} and R_{14} together with the nitrogen to which they are bound are C_4 C_9 heterocycloalkyl, heteroaryl, polyheteroaryl, non-aromatic polyheterocycle or mixed aryl and non-aryl polyheterocycle;
- R_{15} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{16} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, polyheteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{17} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, aromatic polycycle, heteroaryl, arylalkyl, heteroarylalkyl, polyheteroaryl and $NR_{13}R_{14}$; m is an integer selected from 0 to 6; and
- Z is selected from O, NR₁₃, S and S(O);
- or a pharmaceutically acceptable salt thereof.

- 34. A pharmaceutical composition of claim 33 wherein the compound of formula I is selected from the group consisting of N-hydroxy-3-[4-[[(2-hydroxyethyl)]2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, N-hydroxy-3-[4-[[[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide and N-hydroxy-3-[4-[[[2-(2-methyl-1*H*-indol-3-yl)ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 35. A pharmaceutical composition of claim 34 wherein the compound of formula I is N-hydroxy-3-[4-[[(2-hydroxyethyl)]2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 36. A pharmaceutical composition of claim 33 wherein the compound of formula I is N-hydroxy-3-[4-[[[2-(benzofur-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 37. A method for treating a proliferative disorder in a mammal which comprises administering to said mammal a compound of the formula I

HO
$$R_1$$
 R_2 R_3 R_4 R_5 R_5 R_5

wherein

R₁ is H, halo, or a straight chain C₁-C₆ alkyl;

 R_2 is selected from H, C_1 - C_{10} alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, C_4 – C_9 heterocycloalkylalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH_2) $_n$ C(O) R_6 , -(CH_2) $_n$ OC(O) R_8 , amino acyl, HON-C(O)-CH=C(R_1)-aryl-alkyl- and -(CH_2) $_n$ R $_7$;

R₃ and R₄ are the same or different and independently H, C₁-C₆ alkyl, acyl or acylamino, or R₃ and R₄ together with the carbon to which they are bound represent

- C=O, C=S, or C=NR₈, or R₂ together with the nitrogen to which it is bound and R₃ together with the carbon to which it is bound can form a $C_4 C_9$ heterocycloalkyl, a heteroaryl, a polyheteroaryl, a non-aromatic polyheterocycle, or a mixed aryl and non-aryl polyheterocycle ring;
- R₅ is selected from H, C₁-C₈ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, acyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, aromatic polycycle, non-aromatic polycycle, mixed aryl and non-aryl polycycle, polyheteroaryl, non-aromatic polyheterocycle, and mixed aryl and non-aryl polyheterocycle;
- n_1 , n_2 and n_3 are the same or different and independently selected from 0-6, when n_1 is 1-6, each carbon atom can be optionally and independently substituted with R_3 and/or R_4 ;
- X and Y are the same or different and independently selected from H, halo, C₁-C₄ alkyl, NO₂, C(O)R₁, OR₃, SR₃, CN, and NR₁₀R₁;
- R₆ is selected from H, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, OR₁₂, and NR₁₃R₁₄;
- R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{16}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_6$;
- R_8 is selected from H, OR_{15} , $NR_{13}R_{14}$, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;
- R₉ is selected from C₁ C₄ alkyl and C(O)-alkyl;
- R_{10} and R_{11} are the same or different and independently selected from H, C_1 - C_4 alkyl, and -C(O)-alkyl;
- R_{12} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, C_4 C_9 heterocycloalkylalkyl, aryl, mixed aryl and non-aryl polycycle, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, amino acyl, or R_{13} and R_{14} together with the nitrogen to which they are bound are C_4 C_9 heterocycloalkyl, heteroaryl, polyheteroaryl, non-aromatic polyheterocycle or mixed aryl and non-aryl polyheterocycle;
- R_{15} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_mZR_{12}$;
- R_{16} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, polyheteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m$ Z R_{12} ;

 R_{17} is selected from C_1 - C_6 alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkyl, aryl, aromatic polycycle, heteroaryl, arylalkyl, heteroarylalkyl, polyheteroaryl and $NR_{13}R_{14}$; m is an integer selected from 0 to 6; and

Z is selected from O, NR₁₃, S and S(O); or a pharmaceutically acceptable salt thereof.

- 38. A method of claim 37 wherein the compound of formula I is selected from the group consisting of N-hydroxy-3-[4-[[(2-hydroxyethyl)[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, N-hydroxy-3-[4-[[[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-propenamide and N-hydroxy-3-[4-[[[2-(2-methyl-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide, or a pharmaceutically acceptable salt thereof.
- 39. A method for regulating p21 promoter which comprises introducing a compound of the formula I

HO
$$R_1$$
 R_2 R_3 R_4 R_5 R_5 R_5

wherein

R₁ is H, halo, or a straight chain C₁-C₆ alkyl;

 R_2 is selected from H, C_1 - C_{10} alkyl, C_4 – C_9 cycloalkyl, C_4 – C_9 heterocycloalkylalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, -(CH₂)_nC(O)R₈, -(CH₂)_nOC(O)R₈, amino acyl, HON-C(O)-CH=C(R₁)-aryl-alkyl- and -(CH₂)_nR₇;

R₃ and R₄ are the same or different and independently H, C₁-C₈ alkyl, acyl or acylamino, or R₃ and R₄ together with the carbon to which they are bound represent C=O, C=S, or C=NR₈, or R₂ together with the nitrogen to which it is bound and R₃ together with the carbon to which it is bound can form a C₄ - C₉ heterocycloalkyl, a heteroaryl, a polyheteroaryl, a non-aromatic polyheterocycle, or a mixed aryl and non-aryl polyheterocycle ring;

- R₅ is selected from H, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₆ heterocycloalkyl, acyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, aromatic polycycle, non-aromatic polycycle, mixed aryl and non-aryl polycycle, polyheteroaryl, non-aromatic polyheterocycle, and mixed aryl and non-aryl polyheterocycle;
- n_1 , n_1 , n_2 and n_3 are the same or different and independently selected from 0 6, when n_1 is 1-6, each carbon atom can be optionally and independently substituted with R_3 and/or R_4 ;
- X and Y are the same or different and independently selected from H, halo, C₁-C₄ alkyl, NO₂, C(O)R₁, OR₉, SR₉, CN, and NR₁₀R₁₁;
- R_6 is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, cycloalkylalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, OR_{12} , and $NR_{13}R_{14}$;
- R_7 is selected from OR_{15} , SR_{15} , $S(O)R_{16}$, SO_2R_{17} , $NR_{13}R_{14}$, and $NR_{12}SO_2R_6$;
- R₈ is selected from H, OR₁₅, NR₁₃R₁₄, C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, aryl, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_9 is selected from $C_1 C_4$ alkyl and C(O)-alkyl;
- R₁₀ and R₁₁ are the same or different and independently selected from H, C₁-C₄ alkyl, and -C(O)-alkyl;
- R_{12} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, C_4 C_9 heterocycloalkylalkyl, aryl, mixed aryl and non-aryl polycycle, heteroaryl, arylalkyl, and heteroarylalkyl;
- R_{13} and R_{14} are the same or different and independently selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl, amino acyl, or R_{13} and R_{14} together with the nitrogen to which they are bound are C_4 C_9 heterocycloalkyl, heteroaryl, polyheteroaryl, non-aromatic polyheterocycle or mixed aryl and non-aryl polyheterocycle;
- R_{15} is selected from H, C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R_{16} is selected from C_1 - C_6 alkyl, C_4 C_9 cycloalkyl, C_4 C_9 heterocycloalkyl, aryl, heteroaryl, polyheteroaryl, arylalkyl, heteroarylalkyl and $(CH_2)_m ZR_{12}$;
- R₁₇ is selected from C₁-C₆ alkyl, C₄ C₉ cycloalkyl, C₄ C₉ heterocycloalkyl, aryl, aromatic polycycle, heteroaryl, arylalkyl, heteroarylalkyl, polyheteroaryl and NR₁₃R₁₄; m is an integer selected from 0 to 6; and
- Z is selected from O, NR₁₃, S and S(O);
- or a pharmaceutically acceptable salt thereof,

into the environment of a mammalian cell.

40. A method of claim 39 wherein the compound of formula I is selected from the group consisting of N-hydroxy-3-[4-[[(2-hydroxyethyl)[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide, N-hydroxy-3-[4-[[[2-(1H-indol-3-yl)ethyl]-amino]methyl]phenyl]-2E-2-propenamide and N-hydroxy-3-[4-[[[2-(2-methyl-1*H*-indol-3-yl)-ethyl]-amino]methyl]phenyl]-2*E*-2-propenamide, or a pharmaceutically acceptable salt thereof.

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- (71) Applicant (for all designated States except AT, US): NO-VARTIS AG [CH/CH]; Lichtstrasse 35, CH-4056 Basel
- (71) Applicant (for AT only): NOVARTIS-ERFINDUNGEN VERWALTUNGSGESELLSCHAFT M.B.H. [AT/AT]; Brunner Strasse 59, A-1230 Vienna (AT).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): BAIR, Kenneth, Walter [US/US]; 95 Melrose Road, Mountain Lakes, NJ 07046 (US). GREEN, Michael, A. [US/US]; 2180 Biddle Lane, Easton, PA 18040 (US). PEREZ, Lawrence, B. [US/US]; 12 Windsor Place, Hackettstown, NJ 07840 (US). REMISZEWSKI, Stacy, W. [US/US]; 147 Honeysuckle Drive, Washington Township, NJ 07676 (US). SAMBUCETTI, Lidia [US/US]; 32 Lone Mountain Court, Pacifica, CA 94044 (US). VERSACE, Richard,

William [US/US]; 69 Townsend Road, Wanaque, NJ 07465 (US). SHARMA, Sushil, Kumar [US/US]; 9 Bakley Terrace, West Orange, NJ 07052 (US).

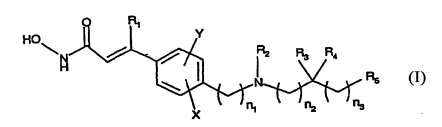
- (74) Agent: BECKER, Konrad; Novartis AG, Corporate Intellectual Property, Patent & Trademark Department, CH-4002 Basel (CH).
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(54) Title: HYDROXAMATE DERIVATIVES USEFUL AS DEACETYLASE INHIBITORS



(57) Abstract: The present invention provides hydroxamate compounds of formula (I) which are deacetylase inhibitors. The compounds are suitable for pharmaceutical compositions having anti-proliferative properties.

INTERNATIONAL SEARCH REPORT

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PCT/EP 01/10037 A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C07D209/16 C07C259/06 C07D417/12 C07D403/12 C07D471/04 A61K31/4045 A61K31/16 C07D519/00 C07D295/02 A61P35/00 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) CO7D CO7C A61K A61P Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, CHEM ABS Data, BIOSIS, WPI Data, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category ° Citation of document, with Indication, where appropriate, of the relevant passages X WO 98 55449 A (QUEENSLAND INST MED RES 1-40 ;FAIRLIE DAVID (AU); PARSONS PETER G (AU);) 10 December 1998 (1998-12-10) claims 4, 15 formula Vg, 16, 25-26; p. 6, 1. 29-33; p. 13-p. 14, 1. 11, p. 45 compound 5, p. 47 3rd structure, p. 48 MW2796, p. 65 table 1 X WO 95 31977 A (SLOAN KETTERING INST CANCER 1 - 40:UNIV COLUMBIA (US)) 30 November 1995 (1995-11-30) claims 40, 42, 55-56; p. 23, 1. 19-1. 35; p. 74 compound 77 and tables 2-3 Χ Patent family members are listed in annex. X Further documents are listed in the continuation of box C. ° Special categories of cited documents: *T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the International "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled *O* document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed in the art. "&" document member of the same patent family Date of the actual completion of the International search Date of mailing of the international search report 30 January 2002 07/03/2002

Authorized officer

Rivat, C

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentiaan 2 NL – 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016

INTERNATIONAL SEARCH REPORT

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PCT/EP 01/10037

	PC1/EP 01/1003				
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	Individual de deservice			
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.			
P,X	WO 01 38322 A (METHYLGENE INC) 31 May 2001 (2001-05-31) claims 26-27,29-31,37,41; compounds 36, 87-91,93-97,99-106, 108-115,120,145-147,170; table 5	1-40			
P,A	M. YOSHIDA ET AL.: "Histone deacetylase as a new target for cancer chemotherapy" CANCER CHEMOTHER. PHARMACOL., vol. 48, no. Suppl. 1, 1 July 2001 (2001-07-01), pages S20-S26, XP002188621 the whole document	1-40			
	~	. ~			

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INTERNATIONAL SEARCH REPORT

infc.....on on patent family members

Internatio Application No
PCT/EP 01/10037

Patent document cited in search report		Publication date		Patent family member(s)		Publication date
WO 9855449	Α	10-12-1998	AU	7751698		21-12-1998
			WO	9855449		10-12-1998
			EP	0988280	A1	29-03-2000
WO 9531977	A	30-11-1995	US	5700811	Α	23-12-1997
			ΑU	692561	B2	11-06-1998
			ΑU	2647495	A	18-12-1995
			CA	2190765	A1	30-11-1995
			ΕP	0760657	A1	12-03-1997
			WO	9531977	A1	30-11-1995
WO 0138322	A	31-05-2001	AU	1876801	Α	04-06-2001
			WO	0138322	Δ1	31-05-2001

Form PCT/ISA/210 (patent family annex) (July 1992)